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Title

High-level language processing regions are not engaged in action observation or imitation

Abbreviated title

Action response in language regions

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Author contributions

E.F. and B.P. designed research, analyzed data, and wrote the manuscript; C.H. helped design, collect data for, and analyze data for Experiments 3a and 3b; K.K. and E.D. helped design and collect data for Experiment 1 and provided comments on the manuscript.

Abstract (234 words)

A set of left frontal, temporal, and parietal brain regions respond robustly during language comprehension and production (e.g., Fedorenko et al. 2010; Menenti et al. 2011). These regions have been further shown to be selective for language relative to other cognitive processes, including arithmetic, aspects of executive function, and music perception (e.g., Fedorenko et al. 2011; Monti et al. 2012). However, one claim about overlap between language and non-linguistic cognition remains prominent. In particular, some have argued that language processing shares computational demands with action observation and/or execution (e.g., Rizzolatti and Arbib 1998; Koechlin and Jubault 2006; Tettamanti and Weniger 2006). However, the evidence for these claims is indirect, based on observing activation for language and action tasks within the same broad anatomical areas (e.g., on the lateral surface of the left frontal lobe). To test whether language indeed shares machinery with action observation/execution, we examined the responses of language brain regions, defined functionally in each individual participant (Fedorenko et al. 2010), to action observation (Experiments 1, 2, 3a) and action imitation (Experiment 3b). With the exception of the language region in the angular gyrus, all language regions, including those in the inferior frontal gyrus (within “Broca’s area”), showed little or no response during action observation/imitation. These results add to the growing body of literature suggesting that high-level language regions are highly selective for language processing (see Fedorenko and Varley 2016 for a review).

54 **New & Noteworthy** (75 words)

55 Many have argued for overlap in the machinery used to interpret language and others'
56 actions, either because action observation was a precursor to linguistic communication or
57 because both require interpreting hierarchically-structured stimuli. However, existing
58 evidence is indirect, relying on group analyses or reverse inference. We examined
59 responses to action observation in language regions defined functionally in individual
60 participants and found no response. Thus, language comprehension and action
61 observation recruit distinct circuits in the modern brain.

62

63 **Introduction**

64 Although brain regions that support high-level language processing have been shown to
65 be selective for language over various non-linguistic cognitive processes (e.g., Fedorenko
66 and Varley 2016), the idea of overlap between language processing and action
67 observation and/or execution remains prominent in the literature. Two lines of theorizing
68 have been used to argue for this overlap. The first stemmed from the discovery of mirror
69 neurons in the prefrontal cortex of rhesus macaques. These neurons fire both when a
70 monkey performs an action and when it observes the action performed (Rizzolatti et al.
71 1988). Rizzolatti & Arbib (1998; Arbib 2005, 2010; see also Petrides & Pandya 2009;
72 Corballis 2010) speculated that in our primate ancestors, mirror neurons were critical for
73 understanding one another's actions – a core component of social cognition. They argued
74 that, over time, basic manual actions grew more abstract, and eventually became signs,
75 which, in turn, became mouth movements/vocalizations. Thus, manual actions are argued
76 to be a fundamental precursor to linguistic communication, and action understanding and
77 language comprehension should share a common neural substrate because they share a
78 common functional ancestor.

79 Although the general idea that language arose from gesture finds substantial
80 support (e.g., Tomasello 2008; Corballis 2003; cf. Slocombe 2015), the role of mirror
81 neurons in the evolution of language remains debated (e.g., Hickok 2009). The existence
82 of brain cells / regions with properties of the macaque mirror neuron system in humans is
83 supported by a number of studies (e.g., Mukamel et al. 2010; see Molenberghs et al. 2012
84 for a meta-analysis) but has not gone unchallenged (e.g., Dinstein et al. 2007; Lingnau et
85 al. 2009). Regardless of these controversies, however, given the prominence of the

gesture-based hypothesis of language evolution, it seems important to test whether any parts of the language network in the modern human brain respond to action observation/execution.

The second general line of reasoning is that both the language system and the action observation system (possibly restricted to biological actions; e.g., Clerget et al. 2009; Fazio et al. 2009) rely on an amodal mechanism that recognizes and produces hierarchical structure (e.g., Fiebach and Schuboltz 2006; Koechlin and Jubault 2006; Tettamanti and Weniger 2006). This mechanism has been argued to reside in the inferior frontal gyrus (IFG), in or around “Broca’s area” (we use quotations because the definition of this brain region in the literature is extremely variable, and the term has been argued by some to no longer be meaningful as a result; Tremblay & Dick, 2016). However, the evidence for overlap between language and action observation in the IFG is problematic because the IFG is among the most structurally (e.g., Amunts et al. 2010) and functionally (e.g., Fedorenko et al. 2012a) heterogeneous brain regions. Further, lateral frontal lobes are characterized by high inter-individual variability (e.g., Amunts et al. 1999; Tomaiuolo et al. 1999; Juch et al. 2005). Thus, activation overlap between language and action observation in a traditional fMRI group analysis (e.g., Higuchi et al. 2009), where activations are averaged across individuals, can be misleading (e.g., Nieto-Castañon and Fedorenko 2012), particularly in the aforementioned regions.

Further, some prior studies did not even include a direct within-experiment comparison between a language and an action task (e.g., Binkofsky et al. 2000; Meister and Iacoboni 2007; Clerget et al. 2009) and relied solely on the fallacious reverse inference (Poldrack 2006, 2011) to interpret the frontal activations for action tasks. This

approach is especially problematic in this case because frontal lobes, including “Broca’s area” itself (Fedorenko et al. 2012a), contain both i) language-selective regions, and ii) highly domain-general ones that belong to the fronto-parietal multiple demand (MD) network (e.g., Duncan 2010) and are driven by diverse cognitive demands (e.g., Duncan & Owen 2000; Fedorenko et al. 2013). Thus, interpreting frontal activations for an action observation task as reflecting the recruitment of the language system is not justified. Similarly, although many aphasic patients with frontal lesions exhibit deficits in action observation/execution (e.g., Kimura 1977; Kimura et al. 1976; Papagno et al., 1993; Saygin et al. 2004), these patients’ lesions are often extensive and plausibly affect two or more functionally distinct regions (cf. Sirugu et al. 1998). Thus, arguing for overlap in mechanisms that support language processing and action observation based on such data is also not warranted.

To test – in the most direct way – whether action observation/execution relies on some of the same neural mechanisms as high-level language processing, we examined responses to action observation and imitation in the language regions functionally defined in each individual. This analytic approach circumvents the problem of high inter-individual variability in the precise locations of functional regions (e.g., Fischl et al. 2008; Frost and Goebel 2011; Tahmasebi et al. 2011) and thus stands a chance to conclusively answer the question about whether language regions support some aspects of action observation. It is worth noting that this question is conceptually distinct from the question that is at the core of the embodiment debate (see Leshinskaya & Caramazza 2016 for a recent review): namely, whether concepts are “grounded” in sensory-motor systems. We elaborate further on the relationship between these questions in the

Discussion.

Materials and Methods

The general approach adopted here across the four experiments is as follows: first, we identify the language network in each participant individually using a functional localizer task based on a broad contrast between the reading of sentences vs. sequences of nonwords (Fedorenko et al. 2010). Then, we examine the engagement of these language-responsive voxels in action observation/imitation across several paradigms. This approach has been previously shown to yield higher sensitivity and functional resolution than traditional group-based analyses, as well as more accurate estimates of effect sizes (e.g., Saxe et al. 2006; Nieto-Castañon and Fedorenko 2012). Further, this approach makes the results directly comparable across the four experiments.

It is worth emphasizing that we here focus on *high-level* language processing regions, i.e., brain regions that support lexico-semantic and combinatorial (semantic and syntactic) processing (e.g., Fedorenko et al. 2012b; Bautista and Wilson 2016; Blank et al. 2016). These regions plausibly underlie our ability to infer meanings from others' linguistic utterances during comprehension as well as to convert our thoughts into linguistic forms during production. This high-level language network is distinct from both lower-level *perceptual* regions that respond selectively to speech, but are not sensitive to the meaningfulness of the speech signal (e.g., Overath et al. 2015; Norman-Haignere et al. 2015) and lower-level *speech articulation* regions that respond robustly when we produce speech sounds, but again are not sensitive to the meaningfulness of the utterance (e.g., Bohland and Guenther, 2006; Flinker et al. 2015; Basilakos et al. 2017).

Thus, our main conclusions pertain to the high-level component of the extended language network. We return to this issue in the Results section.

Participants. Participants were recruited from MIT and the surrounding Cambridge/Boston, MA community and were paid for their participation. Eleven participants were tested in Experiment 1, 57 in Experiment 2, 13 in Experiment 3a, and 16 in Experiment 3b. Seven participants were excluded (3 for excessive motion – all in Experiment 3b, 2 for equipment failure, 1 because an incorrect scanner sequence was used, and 1 due to experimenter error), leaving 90 participants for analysis (10 in Experiment 1, 54 in Experiment 2, 13 in Experiment 3a, and 13 in Experiment 3b). (The number of participants in Experiment 2 was so large because this experiment was used across multiple projects, and we decided to include here all the data available.) Due to some overlap in participants across experiments (8 participated in both Experiment 2 and 3a, and 5 participated in both Experiment 2 and 3b), there were 77 unique individuals (age 18-52, mean age 24, 43 females), 68 right-handed (as determined by the Edinburgh handedness inventory, Oldfield 1971, for n=69, or self report). No participants were excluded based on handedness because we would like to generalize our results to the entire population, as opposed to only the right-handed participants (see Willems et al. 2014, for discussion). The nine left-handed participants all had a left-lateralized language network, as determined by the language localizer task described below. To determine lateralization, the number of language-contrast-activated voxels in the right hemisphere at a fixed significance threshold was subtracted from the number of language voxels in the left hemisphere at the same threshold, and the resulting value was divided by the sum of

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language voxels across hemispheres (see Mahowald & Fedorenko 2016 for further details on this method). All were native speakers of English, had normal hearing and vision, and no history of language impairment. The protocol for these studies was submitted to, and approved by, MIT's Committee on the Use of Humans as Experimental Subjects (COUHES). All participants gave written informed consent in accordance with the requirements of this protocol.

Design and procedure common to all four experiments. Each participant completed a language localizer task (Fedorenko et al. 2010) and an action observation/imitation task. 12 participants completed the localizer task in a separate scanning session; the remaining 78 participants performed the localizer and an action experiment in the same session, along with one or two additional tasks for unrelated studies. The entire scanning session lasted for approximately 2 hours. The task used to localize the language network is described in detail in Fedorenko et al. (2010); the materials and scripts are available from the Fedorenko Lab website (<https://evlab.mit.edu/funcloc>). Briefly, we used a reading task contrasting sentences (e.g., THE SPEECH THAT THE POLITICIAN PREPARED WAS TOO LONG FOR THE MEETING) and lists of unconnected, pronounceable nonwords (e.g., LAS TUPING CUSARISTS FICK PRELL PRONT CRE POME VILLPA OLP WORNIST CHO) in a standard blocked design with a counterbalanced order across runs (for timing parameters, see Table 1). The sentences > nonwords contrast targets brain regions that support lexico-semantic and combinatorial (semantic and syntactic) processing. Stimuli were presented one word/nonword at a time. For 10 participants (in Experiment 1), each trial ended with a memory probe and they had to

indicate, via a button press, whether or not that probe had appeared in the preceding sequence of words/nonwords. The remaining participants instead read the materials passively (we included a button-pressing task at the end of each trial, to help participants remain alert). Importantly, this localizer has been shown to generalize across task manipulations: the sentences > nonwords contrast, and similar contrasts between language and a linguistically degraded control condition, robustly activates the fronto-temporal language network regardless of the task, materials, and modality of presentation (e.g., Fedorenko et al. 2010; Fedorenko 2014; Scott et al. 2016).

The action observation tasks included a variety of conditions – including hand actions with (Experiment 1) or without (Experiment 3a) a manipulable object, actions that involve different body parts including hands, but also arms, legs, feet, torso, and head (Experiment 2), face actions (Experiments 2 and 3a), and specifically eye and mouth actions (Experiment 3a); the action imitation task similarly included several conditions (Experiment 3b). We describe each experiment in more detail below.

Experiment 1: Hand action observation

Participants watched short videos where a small non-nameable 3D object was manipulated in some way by a hand, in a blocked design, and performed a simple one-back task designed to draw attention to the action or the object. (We used non-nameable objects to avoid a potential confound of activating the names of common objects, which would likely elicit some response in the language regions, but not due to overlap in computational demands between language understanding and action observation.) In the action condition, participants had to press a button when they saw the same action twice

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in a row, and in the object condition, they watched the same videos but had to press a button when they saw the same object twice in a row. The task manipulation was included in an effort to maximally focus the participants' attention on the actions in the action condition.

Materials

There were 8 possible hand actions (e.g., push forward with back of the fingers, or pick up with an index finger and a thumb) and 8 possible non-nameable objects, resulting in 64 unique stimuli (see Figure 1 for screenshots from sample stimuli). A short video was created for each action/object combination. Each video started with the object sitting on a table, and then the hand entered the frame (always from the same side), performed the action, and exited the frame. Because objects take less time to identify than actions (given that actions unfold over time), some steps were taken to make the conditions comparable in difficulty. First, the videos were edited so that the action started as quickly as possible after the onset of the video (on average, the action took about 250 ms to initiate). Second, objects were grouped into “families” for presentation purposes such that objects within a family were visually similar to one another. Conversely, actions were grouped in a way such that actions within a set were visually dissimilar.

Procedure

Each video (trial) lasted 3 seconds, and trials were grouped into blocks of 8 trials each. Each block was preceded by a 2-second instructions screen telling participants which condition they were about to see. Each run consisted of 16 such experimental blocks (26 seconds each; 8 blocks per condition) and 5 fixation blocks (16 seconds each, placed at the beginning of the run, and after each set of four blocks). Each run thus lasted 496

seconds (8 min 16 sec). Each participant saw either 4 or 5 runs. The order of conditions was counterbalanced across runs and participants.

Experiment 2: Face and body action observation

Participants passively watched silent videos of i) face actions, ii) body actions, iii) driving through natural scenes, iv) moving man-made objects, and v) spatially scrambled versions of these objects in a blocked design (see Pitcher et al., 2011, for a detailed description). For the purposes of the current study, we examined the first two conditions: face actions and body actions. Participants were instructed to watch attentively.

Materials

There were 60 unique stimuli per condition. The videos depicted children moving against a black background. These children performed a variety of actions like dancing, walking, and crawling (see Figure 1 for screenshots from sample stimuli). The face action videos featured a child's face in motion – smiling, laughing, talking, or looking at someone off-camera. The body action videos featured a child's moving body part – hands, arms, legs, feet, torso, or back of the head – but did not include the face.

Procedure

Each trial consisted of a single video that lasted 3 seconds, and trials were grouped into blocks of 6 trials each. Each run consisted of 10 experimental blocks (18 seconds each; 2 blocks per condition) and 3 fixation blocks (18 seconds each), placed at the beginning, middle, and end of the run. Each run thus lasted 234 seconds (3 min 54 sec). Each participant saw between 2 and 4 runs.

270 ***Experiment 3 a/b: Face, eye, mouth, and hand action observation / imitation***

271 Participants watched silent videos of an actress performing face actions, eye actions,
272 mouth actions, and hand actions. Additionally, the experiment included videos where the
273 actress pronounced consonant and vowel sounds, syllables, nonwords, and words in
274 English and German, and sang or hummed nonwords, all in a blocked design. For the
275 purposes of the current study, we examined the first four conditions: face actions, eye
276 actions, mouth actions, and hand actions. In the observation version of the experiment
277 (Experiment 3a), participants were asked to just watch attentively, and in the imitation
278 version (Experiment 3b), a different set of participants were instructed to imitate each
279 action while keeping their head as still as possible.

280 ***Materials***

281 There were 8 unique stimuli per condition. The videos depicted a female actress against a
282 grey background. In the face, eye, and mouth action conditions, she was sitting facing the
283 camera, with the frame going from just below her shoulders to just above the top of her
284 head. Each video started and ended with the actress looking at the camera, with a neutral
285 expression. The face condition included actions like looking surprised or making a “fish”
286 face (see Figure 1 for screenshots from sample stimuli); the eye condition included
287 actions like moving the eyes up or to the lower left; and the mouth condition included
288 actions like touching the upper teeth with the tongue or pursing the lips to blow air out. In
289 the hand action condition, the hand rested on a wooden table, with the frame covering the
290 hand and a portion of the forearm. Each video started and ended with the hand resting on
291 the table. The hand condition included actions like pulling in the fingers or tapping a
292 finger or multiple fingers on the table.

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293 *Procedure – Experiment 3a (observation)*

294 Each video (trial) lasted 5 seconds, and trials were grouped into blocks of 3 trials each.
295 Each run consisted of 26 experimental blocks (15 seconds each; 2 blocks for each of
296 thirteen conditions) and 5 fixation blocks (14 seconds each), placed at the beginning and
297 end of each run, as well as after the 7th, 13th, and 20th blocks. Each run thus lasted 460
298 seconds (7 min 30 sec). Each participant saw between 4 and 6 runs.

299 *Procedure – Experiment 3b (imitation)*

300 The procedure was identical to that of Experiment 3a except that each video (trial) lasted
301 8 seconds (5 seconds for the video and 3 seconds for the participant to imitate the action;
302 note that although the videos lasted 5 seconds each, the actual action does not take longer
303 than ~3 seconds). Each run thus lasted 694 seconds (11 min 34 sec). Each participant saw
304 between 3 and 8 runs.

305

306 **fMRI data acquisition and preprocessing.** Structural and functional data were collected
307 on the whole-body 3 Tesla Siemens Trio scanner with a 32-channel head coil at the
308 Athinoula A. Martinos Imaging Center at the McGovern Institute for Brain Research at
309 MIT. T1-weighted structural images were collected in 128 axial slices with 1 mm
310 isotropic voxels (TR = 2530 ms, TE = 3.48 ms). Functional, blood oxygenation level
311 dependent (BOLD) data were acquired using an EPI sequence (with a 90 degree flip
312 angle and using GRAPPA with an acceleration factor of 2), with the following
313 acquisition parameters: thirty-one 4 mm thick near-axial slices, acquired in an interleaved
314 order with a 10% distance factor, 2.1 mm x 2.1 mm in-plane resolution; field of view of
315 200 mm in the phase encoding anterior to posterior (A > P) direction; matrix size of 96

mm x 96 mm; TR of 2000 ms; and TE of 30 ms. Prospective acquisition correction (Thesen et al. 2000) was used to adjust the positions of the gradients based on the participant's motion from the previous TR. The first 10s of each run (before the start of presentation of the stimuli) were excluded to allow for steady-state magnetization.

MRI data were analyzed using SPM5 and custom MATLAB and shell scripts. Each participant's data were motion corrected, normalized into a common brain space (MNI) and resampled into 2 mm isotropic voxels. The data were smoothed with a 4mm Gaussian filter and high-pass filtered (at 200s). All task effects were estimated using a General Linear Model (GLM) in which each experimental condition was modeled with a boxcar function convolved with the canonical hemodynamic response function (HRF).

Definition of group-constrained, subject-specific fROIs. The critical analyses were restricted to individually defined language fROIs (functional regions of interest). These fROIs were defined using the Group-constrained Subject-Specific (GSS) approach (Fedorenko et al. 2010; Julian et al. 2012) where a set of spatial parcels (binary masks that correspond to locations where activation has been previously observed for the relevant localizer contrast) is combined with each individual subject's localizer activation map, to constrain the definition of individual fROIs. The parcels are sufficiently large to encompass the extent of variability in the locations of individual activations. For the critical language fROIs, we used a set of six parcels derived from a group-level probabilistic activation overlap map for the sentences > nonwords contrast in 220 participants. These parcels (Figure 2) included three regions in the left frontal cortex: two in the left inferior frontal gyrus (LIFG, LIFGorb), and one in the left middle frontal gyrus

(LMFG), two in the left temporal lobe (LAntTemp and LPostTemp), and one extending into the angular gyrus (LAngG). These parcels are similar to the ones originally reported in Fedorenko et al. (2010) based on a probabilistic activation overlap map from 25 participants, except that the two anterior temporal parcels were grouped together (the original LAntTemp merged with LMidAntTemp), and the two posterior temporal parcels were grouped together (the original LMidPostTemp merged with LPostTemp). The parcels are available for download from <https://evlab.mit.edu/funcloc>.

Within each parcel, we selected the top 10% most responsive voxels, based on the t values for the sentences > nonwords contrast (see e.g., Figure 1 in Blank et al. 2014; or Figure 1 in Mahowald and Fedorenko 2016, for sample fROIs). Statistical tests were performed on these values.

In addition to the language fROIs, a set of control fROIs was defined in the participants in Experiments 2, 3a, and 3b. In particular, we used 18 anatomical parcels across the two hemispheres (Tzourio-Mazoyer et al. 2002) covering frontal and parietal brain areas that belong to the so-called multiple demand (MD) network (Duncan 2010, 2013). This network has been linked to executive demands across domains (e.g., Duncan and Owen 2000; Fedorenko et al. 2013; Hugdahl et al. 2015), but parts of this network have also been implicated in the processing of actions (e.g., Culham and Valer 2006; Gallivan and Culham 2015; Biagi et al. 2015; Caspers et al. 2010). We thus expected some of these regions to respond to action observation and/or imitation. In particular, we focused on a subset of 6 parcels (although the results were corrected for the total number of regions that comprise this network, i.e., 18): the bilateral IFGop and PrecG fROIs in the frontal cortex because those lie in close proximity to the language fROIs, and the

bilateral SupPar fROIs in the parietal cortex because these regions have been implicated in action observation in prior work (e.g., Johnson-Frey et al., 2005).

To define individual MD fROIs, we used a spatial working memory task where participants keep track of locations within a grid (this MD localizer task was not included in Experiment 1, hence this analysis could not be performed for those participants). The task is described in detail in Fedorenko et al. (2013; see also Blank et al. 2014). Briefly, on each trial, participants saw a 3×4 grid and kept track of eight (hard version) or four (easy version) locations that were sequentially flashed two at a time or one at a time, respectively. Then, participants indicated their memory for these locations in a two-alternative, forced-choice paradigm via a button press. Feedback was provided after every trial. Hard and easy conditions were presented in a standard blocked design (4 trials in a 32s block, 6 blocks per condition per run) with a counterbalanced order across runs. Each run included 4 blocks of fixation (16s each) and lasted a total of 448s. Within each anatomical parcel, we selected the top 10% most responsive voxels, based on the *t* values for the hard > easy spatial working memory contrast. Statistical tests were performed on these values.

Finally, for some additional analyses reported in the Discussion, we examined i) brain regions in the auditory cortex that support speech perception, and ii) brain regions in the premotor cortex that support speech articulation. For the former, we used the following anatomical parcels from the FSL atlas (Desikan et al. 2006): bilateral planum polare (PP), planum temporale (PT), anterior superior temporal gyrus (ASTG), and posterior superior temporal gyrus (PSTG). To define individual speech-responsive fROIs, these anatomical parcels were masked with activation maps for a contrast between

listening to nonwords and observing hand actions (in Experiment 3a). The responses were then extracted to nonwords, and the four action observation conditions. To estimate the responses to the nonwords and hand action observation conditions, an across-runs cross-validation procedure was used so that the data to define the fROIs and estimate their responses were independent (e.g., Kriegeskorte et al. 2011). In particular, all but one run were used to define the fROIs and the responses were estimated in the left-out run; this procedure was repeated leaving out each run in turn; the response estimates were then averaged across runs to derive a single estimate per condition per fROI. This procedure allows all of the data to be used while maintaining the independence between the data used to define the fROIs and the data used to examine their responses (e.g., Nieto-Castañón & Fedorenko 2012).

For the articulation regions, we used functional parcels derived from a group-level probabilistic activation overlap map for the contrast between the production of difficult-to-articulate nonwords and fixation in 20 participants, as reported in Basilakos et al. (2017). We focused on the regions in the premotor cortices bilaterally: a region in the left precentral gyrus, and two regions in the right precentral gyrus (see Figure 3 in Basilakos et al. 2017). To define individual articulation-responsive fROIs, these parcels were masked with activation maps for a contrast between imitating nonwords (repeating the nonword produced by the actress) and imitating hand actions (in Experiment 3b). The responses were then extracted to nonwords, and the four action imitation conditions. As with the analyses of the speech-responsive regions, to estimate the responses to the nonwords and hand action imitation conditions, an across-runs cross-validation procedure was used so that the data to define the fROIs and estimate their responses were

independent (e.g., Kriegeskorte et al. 2011).

Analyses. In the critical analyses that examined the responses of the language fROIs to the different action observation / imitation conditions, we used two-tailed t -tests to compare the responses to each action condition against i) the low-level fixation baseline, ii) nonword processing, which serves as the control condition in the language localizer, and iii) sentence comprehension. The resulting p values were corrected for the number of language fROIs within each experiment (i.e., 6), using the False Discovery Rate correction (Benjamini and Yekutieli 2001). If language comprehension and action observation / imitation share computational demands, then the action conditions should elicit a response that is as strong as the sentence comprehension condition, or, at least, reliably stronger than the nonword processing condition.

Results

Behavioral data

Overt behavioral responses were only collected in Experiment 1, where participants watched videos and performed a one-back task on the action or the object in the video, as described in Methods. Accuracies were high in both conditions, but slightly and reliably higher for the actions condition than the objects condition (94.9% and 87.5%, respectively; two-tailed $t(9) = 3.18$, $p < 0.05$). Further, as expected (given that actions take time to unfold), participants were faster in the objects condition than the actions condition (1.37s vs. 1.71s; two-tailed $t(9) = 6.05$, $p \leq 0.0005$).

Validation of the language fROIs

Replicating previous work (Fedorenko et al. 2010; Fedorenko et al. 2011), the sentences > nonwords effect was highly reliable in each of six fROIs both i) across the entire set of participants ($ts(76) > 10$, $ps < 0.0001$), and ii) in each experiment individually (Experiment 1: $ts(9) > 4.43$, $ps < 0.001$, Experiment 2: $ts(53) > 8.39$, $ps < 0.0001$, Experiment 3a: $ts(12) > 3.68$, $ps < 0.005$, and Experiment 3b: $ts(12) > 4.01$, $ps < 0.001$). Here, and in validating the MD fROIs, an across-runs cross-validation procedure, described above, was used so that data used to define the fROIs were independent of the data used to estimate the responses.

Responses of the language fROIs to the action conditions

The results are reported in Table 2 and Figure 2. Across experiments, none of the language regions responded strongly and consistently to action observation or imitation. In most fROIs, the action conditions failed to elicit a response above the fixation baseline (except for Experiment 2, where both conditions elicited small but reliable above-baseline responses in all language fROIs). Further, the response to the action observation/imitation condition did not significantly differ from the nonword condition, with the exception of the AngG fROI, which responded more strongly to some action observation conditions than the nonword condition. Finally, again with the exception of the AngG fROI, the response to the action observation/imitation condition was reliably (in almost all cases, and always numerically) below that elicited by sentence comprehension.

Experiment 1. When participants watched videos of a hand performing simple manipulations of an object, there was no above-baseline response in any of the language fROIs, regardless of whether participants were asked to focus on the objects ($ts(9) < 1.5$,

n.s.) or actions ($ts(9) < 1.6$, n.s.). Further, neither of the action conditions elicited a response that was reliably greater than the nonword condition, whereas the sentence condition elicited a reliably greater response than either of the two action conditions ($ts(9) > 2.75$, $ps < 0.05$).

Experiment 2. In this experiment, every language fROI showed a reliably above-baseline response to both the face action observation condition ($ts(53) > 2.11$, $ps < 0.05$) and the body action observation condition ($ts(53) > 2.86$, $ps < 0.01$). However, in all fROIs except for the AngG fROI, this response was i) not reliably higher than that elicited by the nonword condition ($ts(53) < 1.67$, $ps > 0.16$), and ii) reliably lower than that elicited by the sentence condition ($ts(53) > 5.72$, $ps < 0.0001$). In the AngG language fROI, both action observation conditions elicited a response that was reliably stronger than that elicited by the nonword condition and that did not differ from that elicited by the sentence condition. We come back to the AngG fROI in the Discussion.

Experiment 3a. Similar to Experiment 1, there was no above-baseline response in the language fROIs to any of the four conditions, with the exception of the AngG fROI and the MFG fROI, which showed reliably above-baseline responses to hand action observation ($ts(12) > 2.82$, $ps < 0.05$), but only the AngG fROI responded reliably more strongly to hand action observation (and mouth action observation) than to nonwords ($ts(12) > 3.67$, $ps < 0.05$); in all other fROIs none of the action observation conditions produced a stronger response than nonwords. Finally, in all language fROIs, except for the AngG fROI, the sentence condition elicited a reliably greater response than each of the four action observation conditions ($ts(12) > 3.30$, $ps < 0.01$). In the AngG fROI, the response to the action observation conditions did not reliably differ in magnitude from

the sentence condition.

Experiment 3b. In this experiment, where participants observed and imitated different kinds of actions, there was no above-baseline responses except for the MFG fROI, which responded reliably above baseline to the eye, mouth, and hand action conditions ($t(12) > 2.23$, $ps < 0.05$), and marginally to the face action condition ($t(12) = 3.09$, $p = 0.056$). However, these responses did not significantly differ from the response elicited by the nonword condition (see Fedorenko et al. 2011, for a similar pattern of results with other non-linguistic tasks). Further, the sentence condition elicited a reliably or marginally greater response than each of the four action conditions in all language fROIs, except for the AngG fROI and some frontal fROIs for some of the conditions (see Table 2 for details).

Validation of the control, multiple demand (MD), fROIs

Replicating previous work (Fedorenko et al. 2013; Blank et al. 2014), the hard > easy spatial working memory effect was highly reliable in each of six fROIs across participants with 2 runs ($ts(47) > 7.8$, $ps < 0.0001$). Participants with 1 run only ($n=18$) could not be included in this validation analysis because across-runs cross-validation could not be performed; for those participants, we ensured that MD activations looked as expected based on visual examination of whole-brain activation maps.

Responses of the control, multiple demand (MD), fROIs to the action conditions

Unlike in the language fROIs, all action imitation conditions elicited reliably above-baseline responses in almost all MD fROIs. Similarly, at least some action observation conditions elicited reliable responses. The body action observation condition from Experiment 2, and the eye and hand action observation conditions from Experiment 3a

elicited the strongest responses. Strong responses to eye movement observation and imitation could be related to prior claims about the role of this fronto-parietal system in saccades (e.g., Pierrot-Deseilligny et al. 2004).

Responses of speech perception and articulation regions to the action conditions

As discussed at the beginning of the Methods section, we have here focused on high-level language processing regions (e.g., Fedorenko et al. 2010), which plausibly store our linguistic knowledge that we use to both interpret and generate meaningful utterances (e.g., Menenti et al. 2011). These regions are distinct from lower-level speech perception regions (e.g., Overath et al. 2015; Norman-Haignere et al. 2015) and from speech articulation regions (e.g., Bohland and Guenther 2006; Flinker et al. 2015; Basilakos et al. 2017). Might some of this perceptual or motor speech machinery overlap with action observation or imitation? Based on the available evidence, a tentative answer appears to be ‘no’. In particular, the superior temporal regions that respond robustly to speech show some response during speech articulation (e.g., Hickok et al. 2009; Basilakos et al. 2017), but respond very little when participants produce even actions that involve speech articulators, i.e., non-speech oral-motor movements (Basilakos et al. 2017). To shed further light on this question, we performed an additional analysis on data from Experiment 3a. We used a contrast between listening to nonwords and hand action observation to define speech-responsive regions within the superior temporal cortex, and then examined the responses of those regions to nonwords and hand action observation (in data not used for fROI definition), as well as to face, eye, and mouth action observation conditions. As Figure 4a clearly shows, the four action observation conditions fail to elicit above-baseline responses, suggesting that these regions do not

support action observation.

What about regions that support speech articulation? Basilakos et al. (2017) report quite robust responses to the production of non-speech oral-motor movements in premotor articulation regions. We performed an additional analysis on data from Experiment 3b to examine the responses of those articulation regions to action imitation more broadly. We used a contrast between imitating nonwords (repeating the nonword produced by the actress) and hand actions to define articulation-responsive regions within ventral premotor cortex, and then examined the responses of those regions to nonwords and hand action imitation (in data not used for fROI definition), as well as to face, eye, and mouth action imitation. As Figure 4b shows, the mouth action imitation condition elicits as strong a response as, or a stronger response than, articulation, replicating Basilakos et al. (2017). The face condition (which also includes some mouth movements) also elicits a strong response. However, the hand and eye action imitation conditions elicit much lower responses. This relative selectivity for speech and oral-motor/face actions is in line with the idea that these regions contain a map of our articulatory apparatus (e.g., Bouchard et al. 2013; Guenther 2016), arguing against broad engagement in action imitation, as well as with prior findings of somatotopic organization in the motor areas (e.g., Watkins et al. 2003; Pulvermuller et al. 2006; D'Ausilio et al. 2009; Murakami et al. 2011).

Thus, similar to high-level language processing regions, speech perception regions do not appear to support action observation, and speech articulation regions do not appear to support action imitation.

546 **Discussion**

547 We asked whether any part(s) of the language network – a set of brain regions that
548 support high-level language processing (e.g., Fedorenko et al. 2010; Fedorenko &
549 Thompson-Schill, 2014) – respond to action observation and/or imitation. Neural
550 machinery that supports both language processing and some aspects of action
551 observation/imitation has been postulated based on two distinct ideas. First, inspired by
552 the discovery of mirror neurons in macaques (Rizzolatti et al. 1988), some have argued
553 that manual actions served as a fundamental precursor to linguistic communication in the
554 evolution of our species (e.g., Arbib 2005; but see e.g., Tomasello 2008; Corballis 2003,
555 for arguments for gesture-based origins of language that do not hinge on the mirror-
556 neuron-based theorizing). Second, some have postulated an amodal hierarchical
557 processor in the left frontal cortex (in or near “Broca’s area”) that is hypothesized to
558 support both language processing and action perception/planning (e.g., Tettamanti and
559 Weniger 2006; Fiebach and Schuboltz 2006; Koechlin and Jubault 2006).

560 Across three experiments (77 participants, 90 scanning sessions), we examined
561 neural responses of functionally defined language regions to a broad range of *action*
562 *observation* conditions, including hand actions with (Experiment 1) or without
563 (Experiment 3a) a manipulable object, but also actions that involve the face or face parts
564 (Experiments 2 and 3a) and body parts other than the hands (Experiment 2). In the fourth
565 experiment (13 participants), we further examined responses of language regions to
566 *action imitation*, again involving different face and body parts.

567 The key result is that – with a single exception discussed below – none of the
568 language regions responded strongly and consistently to action observation or imitation.

Action response in language regions

In most language regions, the action conditions did not elicit a response above the fixation baseline, which suggests that the language regions are as active during action observation/imitation as they are when we are looking at a blank screen. Further, in most language regions, the response to the action observation/imitation conditions i) did not significantly differ from the response elicited by the nonword condition (the control condition in the language localizer task), and ii) was reliably lower than the response elicited by the sentence condition. These results suggest that language regions are selective for language processing, in line with earlier work that established selectivity for language relative to arithmetic, executive processing, music perception, and social cognition (e.g., Fedorenko and Varley 2016). This conclusion is also consistent with lesion studies that have reported dissociations between linguistic deficits and deficits in action observation/production (e.g., Sirigu et al. 1998), and with a recent fMRI study that showed that the degree of lateralization for language appears to be unrelated to the degree of lateralization for action observation (Häberling et al. 2016).

The only exception was the language fROI in the angular gyrus. This region responded more strongly to some action observation conditions than to nonwords, and, in some cases, the response to action observation was not significantly lower than the response to sentences. Evidence is accumulating that this region differs functionally from the rest of the language network. In particular, it shows relatively low functional correlations with other language regions during naturalistic cognition (e.g., Blank et al. 2014), including when using dynamic network modeling (Chai et al. 2016), and it shows lower correlations in effect sizes and lateralization (e.g., Mahowald and Fedorenko 2016). It also differs from the other language regions in sensitivity to linguistic and non-

linguistic manipulations. For example, the AngG language fROI was the only region that did not show sensitivity to syntactic complexity (Blank et al. 2016), and it was the only region that did not respond more strongly to sentences than photographs matched for semantic content (Amit et al. 2017). The latter result suggests that the AngG language fROI may respond to visual stimuli in general, as opposed to action observation specifically. However, the precise role of this region in human cognition remains to be discovered. One current hypothesis (formulated not specifically about the language-responsive portion of the angular gyrus, but about the broad anatomical area) is that it is “involved in all aspects of semantic processing” and contributes to “behaviors requiring fluent conceptual combination” (e.g., Binder et al. 2009; cf. Lambon Ralph et al. 2017).

We now touch on four theoretical issues that the current results bear on.

Gestural origins of language

Just because in the modern human brain, language processing and action observation appear to recruit non-overlapping machinery does not imply that our linguistic communication system could not have arisen from the manual modality. In fact, this possibility is still perhaps the most plausible (e.g., Tomasello 2008; Corballis 2003; cf. Slocombe 2015; Shepherd & Freiwald, 2018). However, once humans began to develop an extensive set of vocal communication signals, they plausibly had to allocate some portions of the association cortices – massively expanded in the human brain (e.g., Buckner and Krienen 2013) – to store these form-meaning mappings (see also Häberling et al., 2016). Given the differences between linguistic and non-linguistic communication signals (including both discrete, categorical speech-accompanying gestures and

continuous, mimetic facial expressions and body language) – it is perhaps to be expected that these different forms of communication would recruit distinct cognitive (e.g., Goldin-Meadow and Brentari 2016; McNeill 1992) and neural (e.g., Häberling et al. 2016) machinery given the distinct computational demands they place on the mind and brain. It is worth noting that a few prior neuroimaging studies have argued that gesture processing does recruit the same brain regions as language comprehension (e.g., Villarreal et al. 2008; Xu et al. 2009; Enrici et al. 2011; Andric et al. 2013; Redcay et al. 2016; see Willems & Hagoort 2007; Marstaller & Burianová 2014; Yang et al. 2015, for reviews). However, those studies typically used symbolic gestures, pantomime, or “emblems” (e.g., wave, hold out hand for a shake, etc.). Given that such gestures are clearly associated with particular meanings, their processing may lead to the activation of the corresponding linguistic representations. Thus, the overlap may be explained by the engagement of linguistic resources during the processing of gestures rather than by the shared computational demands like communicative intent or abstract conceptual processing.

Amodal hierarchical processor in “Broca’s area”?

Although our action observation/imitation conditions did not include a manipulation of hierarchical complexity, we would argue that – to the extent that an amodal hierarchical processor exists in the human brain – it does not reside within the high-level language network. We have previously made this argument based on non-overlap between language processing and music perception (e.g., Fedorenko et al. 2011; Fedorenko et al. 2012c; Norman-Haignere et al. 2015). Music is another domain that has been argued to

recruit such an amodal hierarchical processor (e.g., Maess et al. 2001; Koelsch et al. 2002). However, as Fedorenko & Varley (2016) have argued, the most compelling evidence for overlap comes from structure-violation paradigms, and in those paradigms, violations of structure appear to elicit similar responses to those elicited by low-level oddball manipulations (e.g., Corbetta and Shulman 2002) and plausibly arise within the domain-general multiple demand (MD) network due to increased processing effort associated with unexpected events (Duncan 2010, 2013). Similarly, some manipulations of hierarchical complexity in the action domain (e.g., Koechlin and Jubault 2006) plausibly engage parts of the MD network because more complex action plans are associated with greater working memory and cognitive control demands. Although parts of the MD system have been argued to be particularly sensitive to hierarchical demands (e.g., Badre and D'Esposito 2007, 2009; Badre 2008) or to the level of abstractness of the to-be-processed information (e.g., Koechlin et al. 2003; Koechlin and Summerfield 2007), these proposals have not gone unchallenged (e.g., Crittenden and Duncan 2012; Pischedda et al. 2017). Thus, whether an amodal hierarchical processor exists anywhere in the human brain remains an open question, but to the extent that it does, it exists outside the boundaries of the high-level language network.

Possibly similar computations across domains in spite of non-overlapping brain regions

The existence of distinct brain regions that support information processing in different domains – like language vs. action observation vs. action production – does not imply that the basic computations (that operate over those domain-specific representations) are

different. In fact, neural circuits across the cortex share many core properties (e.g., Douglas et al. 1989; Douglas and Martin 2004; Harris and Shepherd 2015), suggesting that the basic computations may be the same or similar across different cortical areas. It is also easy to come up with intuitive-level descriptions of potential parallels between domains. For example, in the domain of language, we have a large store of form-meaning mappings and knowledge about the relationships among them. We can use this knowledge to interpret linguistic signals, and to generate new utterances, by combining these basic building blocks into sequences. In the domain of actions, we may have a similar “vocabulary” of actions for each of our effectors associated with particular contexts of use, and information about how these actions can be combined (e.g., Hommel et al. 2001; Rosenbaum et al. 2001; Schack 2004). And we can refer to this stored knowledge to interpret others’ actions as well as generate our own action sequences as needed for goal-directed behavior (e.g., Flash and Bizzi 2016). As we make progress in developing fleshed-out mechanistic-level hypotheses about what actually goes on when we understand and produce language, or as we perceive and generate motor actions, it is important to keep in mind both that i) the linguistic and action/motor representations appear to be stored in non-overlapping brain areas, but that ii) the computations may be fundamentally similar between these (and possibly other domains of perception, action, and cognition).

(Ir)relevance of the current results to the embodiment debate

As noted in the Introduction, the question investigated here – i.e., whether high-level language processing brain regions are engaged when we observe or produce motor

actions – is distinct from the much-debated question of the *nature of our conceptual representations*. In particular, for many years now, some have advocated an “embodied” view of meanings whereby concepts are “grounded” in sensory-motor modalities (e.g., Barsalou et al. 2003; Tranel et al. 2003; Simmons et al. 2007). Embodiment proposals vary widely in the scope of their claims (see Leshinskaya and Caramazza 2016, for a recent review of the key issues in this debate), from a complete denial of the existence of abstract / amodal conceptual representations (e.g., Barsalou et al. 2003; Barsalou 2008; Pulvermüller and Fadiga 2010; cf. Caramazza et al. 1990) to more moderate positions where abstract representations interact in some way with the sensory/motor ones (e.g., Meteyard et al. 2012). The reason that the work reported here might, on the surface, appear to be relevant to the embodiment debate is that action verbs have received a lot of attention in that literature (e.g., see Bedny and Caramazza 2011, for a review). However, the link is superficial: whether or not sensory and/or motor brain regions are active (to some extent) when we understand the meanings of verbs like “kick” or “punch” (e.g., Hauk et al. 2004) is orthogonal to the question of whether the regions of the language network – that we *know* are engaged when we process word meanings (e.g., Fedorenko et al., 2012b) – play a role in the processing or execution of motor actions. We here show that the answer to the latter question is no.

Before concluding, it is worth noting that, in general, the construct of “actions” is complex and heterogeneous, and different researchers have different notions and scope in mind when they talk about “actions”. The conditions we included in our study have spanned goal-directed/transitive actions (e.g., manipulating an object in Experiment 1)

Action response in language regions

and intransitive ones (e.g., tapping a finger in Experiment 3a), as well as actions performed by different effectors (hand, feet, eyes, mouth, face). There are plausibly other dimensions of actions that affect their representation and processing (e.g., Tarhan & Konkle, 2017): e.g., whether the action is directed toward an animate entity vs. an object, whether the action has communicative intent, whether the action maps onto a linguistic label, etc. Our study leaves open the possibility that actions that have some specific property/-ies may elicit responses in the language cortex. However, to evaluate this possibility, we need clear testable hypotheses that would formally specify the relevant features of actions that may lead to the recruitment of the same machinery as language comprehension.

To conclude, action observation and action imitation do not recruit the left-lateralized high-level language processing network, providing further evidence for the selectivity of this network for language processing (e.g., Fedorenko and Varley 2016). However, this separability is still compatible with hypotheses about the gestural origins of human language (e.g., Tomasello 2008; Corballis 2003). Further, given the general similarity of neural circuits across the neocortex (e.g., Harris and Shepherd 2015), research in the domains of action perception or motor control may inform our understanding of the computations that support language comprehension and production, domains where we don't have the luxury of animal models to richly characterize neural response properties and their interactions.

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Conflict of interest

The authors declare no competing financial interests.

747 **References**

- 748
749 **Amit E, Hoeflin C, Hamzah N, Fedorenko E.** An asymmetrical relationship between
750 verbal and visual thinking: Converging evidence from behavior and fMRI.
751 *NeuroImage* 152: 619-627, 2017.
- 752 **Amunts K, Schleicher A, Bürgel U, Mohlberg H, Uylings HBM, Zilles K.** Broca's
753 Region Revisited: Cytoarchitecture and Intersubject Variability. *J Comp Neurol*
754 412: 319-341, 1999.
- 755 **Amunts K, Lenzen M, Friederici AD, Schleicher A, Morosan P, Palomero-Gallagher**
756 **N, Zilles K.** Broca's Region: Novel Organizational Principles and Multiple
757 Receptor Mapping. *PLoS Biol* 8: e1000489, 2010.
- 758 **Andric M, Solodkin A, Buccino G, Goldin-Meadow S, Rizzolatti G, Small SL.** Brain
759 Function Overlaps When People Observe Emblems, Speech, and Grasping.
760 *Neuropsychologia* 51: 1619-1629, 2013.
- 761 **Arbib MA.** From monkey-like action recognition to human language: An evolutionary
762 framework for neurolinguistics. *Behav Brain Sci* 28: 105-167, 2005.
- 763 **Arbib MA.** Mirror system activity for action and language is embedded in the integration
764 of dorsal and ventral pathways. *Brain Lang* 112: 12-24, 2010.
- 765 **Badre D, D'Esposito M.** Functional Magnetic Resonance Imaging Evidence for a
766 Hierarchical Organization of the Prefrontal Cortex. *J Cognitive Neurosci* 19:
767 2082-2099, 2007.
- 768 **Badre D.** Cognitive control, hierarchy, and the rostro-caudal organization of the frontal
769 lobes. *Trends Cogn Sci* 12: 193-200, 2008.
- 770 **Badre D, D'Esposito M.** Is the rostro-caudal axis of the frontal lobe hierarchical?. *Nat*

- 771 *Neurosci* 10: 659-669, 2009.
- 772 **Barsalou LW, Simmons WK, Barbey AK, Wilson CD.** Grounding conceptual
773 knowledge in modality-specific systems. *Trends Cogn Sci* 7: 84-91, 2003.
- 774 **Barsalou LW.** Grounded cognition. *Annu Rev Psychol* 59: 617-645, 2008.
- 775 **Basilakos A, Smith KG, Fillmore P, Fridriksson J, Fedorenko E.** Functional
776 Characterization of the Human Speech Articulation Network. *Cereb Cortex* 1-15,
777 2017.
- 778 **Bautista A, Wilson SM.** Neural responses to grammatically and lexically degraded
779 speech. *Lang Cogn Neurosci* 31: 567-574, 2016.
- 780 **Bedny M, Caramazza A.** Perception, action, and word meanings in the human brain: the
781 case from action verbs. *Ann NY Acad Sci* 1224: 81-95, 2011.
- 782 **Benjamini Y, Yekutieli D.** The Control of the False Discovery Rate in Multiple Testing
783 under Dependency. *Ann Stat* 29: 1165-1188, 2001.
- 784 **Biagi L, Cioni G, Fogassi L, Guzzetta A, Sgandurra G, Tosetti M.** Action observation
785 network in childhood: a comparative fMRI study with adults. *Dev Sci* 19: 1075-
786 1086, 2016.
- 787 **Binder JR, Desai RH, Graves WW, Conant LL.** Where Is the Semantic System? A
788 Critical Review and Meta-Analysis of 120 Functional Neuroimaging Studies.
789 *Cereb Cortex* 19: 2767-2796, 2009.
- 790 **Binkofski F, Amunts K, Stephan KM, Posse S, Schormann T, Freund HJ, Zilles K,**
791 **Seitz RJ.** Broca's Region Subserves Imagery of Motion: A Combined
792 Cytoarchitectonic and fMRI Study. *Hum Brain Mapp* 11: 273-285, 2000.
- 793 **Blank I, Kanwisher N, Fedorenko E.** A functional dissociation between language and

- 794 multiple-demand systems revealed in patterns of BOLD signal fluctuations. *J*
795 *Neurophysiol* 112: 1105-1118, 2014.
- 796 **Blank I, Balewski Z, Mahowald K, Fedorenko E.** Syntactic processing is distributed
797 across the language system. *NeuroImage* 127: 307-323, 2016.
- 798 **Bohland JW, Guenther FH.** An fMRI investigation of syllable sequence production.
799 *NeuroImage* 32: 821-841, 2006.
- 800 **Bouchard KE, Mesgarani N, Johnson K, Chang EF.** Functional organization of human
801 sensorimotor cortex for speech articulation. *Nature* 495: 327-332, 2013.
802 (Corrigendum. *Nature* 498: 526, 2013)
- 803 **Buckner RL, Krienen FM.** The evolution of distributed association networks in the
804 human brain. *Trends Cogn Sci* 17: 648-665, 2013.
- 805 **Caramazza A, Hillis AE, Rapp BC, Romani C.** The Multiple Semantics Hypothesis:
806 Multiple Confusions?. *Cogn Neuropsychol* 7: 161-189, 1990.
- 807 **Caspers S, Zilles K, Laird AG, Eickhoff SB.** ALE meta-analysis of action observation
808 and imitation in the human brain. *NeuroImage* 50: 1148-1167, 2010.
- 809 **Chai LR, Mattar MG, Blank IA, Fedorenko E, Bassett DS.** Functional Network
810 Dynamics of the Language System. *Cereb Cortex* 1-12, 2016.
- 811 **Clerget E, Winderickx A, Fadiga L, Olivier E.** Role of Broca's area in encoding
812 sequential human actions: a virtual lesion study. *NeuroReport* 20: 1496-1499,
813 2009.
- 814 **Corballis MC.** From hand to mouth: The origins of language. Princeton, NJ: Princeton,
815 2003.
- 816 **Corballis MC.** Mirror neurons and the evolution of language. *Brain Lang* 112: 25-35,

- 817 2010.
- 818 **Corbetta M, Shulman GL.** Control of goal-directed and stimulus-driven attention in the
819 brain. *Nat Neurosci* 3: 201-215, 2002.
- 820 **Crittenden BM, Duncan J.** Task Difficulty Manipulation Reveals Multiple Demand
821 Activity but no Frontal Lobe Hierarchy. *Cereb Cortex* 24: 532-540, 2012.
- 822 **Culham JC, Valyear KF.** Human parietal cortex in action. *Curr Opin Neurobiol* 16:
823 205-212, 2006.
- 824 **D'Ausilio A, Pulvermüller F, Salmas P, Bufalari I, Begliomini C, Fadiga L.** The
825 Motor Somatotopy of Speech Perception. *Curr Biol* 19: 381-385, 2009.
- 826 **Desikan RS, Ségonne F, Fischl B, Quinn BT, Dickerson BC, Blacker D, Buckner RL,**
827 **Dale AM, Maguire RP, Hyman BT, Albert MS, Killiany RJ.** An automated
828 labeling system for subdividing the human cerebral cortex on MRI scans into
829 gyral based regions of interest. *NeuroImage* 31: 968-980, 2006.
- 830 **Dinstein I, Hasson U, Rubin N, Heeger DJ.** Brain Areas Selective for Both Observed
831 and Executed Movements. *J Neurophysiol* 98: 1415-1427, 2007.
- 832 **Douglas RJ, Martin KAC, Whitteridge D.** A Canonical Microcircuit for Neocortex.
833 *Neural Comput* 1: 480-488, 1989.
- 834 **Douglas RJ, Martin KAC.** Neuronal circuits in the neocortex. *Ann Rev Neurosci* 27:
835 419-451, 2004.
- 836 **Duncan J, Owen AM.** Common regions of the human frontal lobe recruited by diverse
837 cognitive demands. *Trends Neurosci* 23: 475-483, 2000.
- 838 **Duncan J.** The multiple-demand (MD) system of the primate brain: mental programs for
839 intelligent behaviour. *Trends Cogn Sci* 14: 172-179, 2010.

- 840 **Duncan J.** The Structure of Cognition: Attentional Episodes in Mind and Brain. *Neuron*
841 80: 35-50, 2013.
- 842 **Enrici I, Adenzato M, Cappa S, Bara BG, Tettamanti M.** Intention Processing in
843 Communication: A Common Brain Network for Language and Gestures. *J*
844 *Cognitive Neurosci* 23: 2415-2431, 2011.
- 845 **Fazio P, Cantagallo A, Craighero L, D'Ausilio AD, Roy AC, Pozzo T, Calzolari F,**
846 **Granieri E, Fadiga L.** Encoding of human action in Broca's area. *Brain* 132:
847 1980-1988, 2009.
- 848 **Fedorenko E, Hsieh PJ, Nieto-Castañón A, Whitfield-Gabrieli S, Kanwisher N.** New
849 Method for fMRI Investigations of Language: Defining ROIs Functionally in
850 Individual Subjects. *J Neurophysiol* 104: 1177-1194, 2010.
- 851 **Fedorenko E, Behr MK, Kanwisher N.** Functional specificity for high-level linguistic
852 processing in the human brain. *P Natl Acad Sci USA* 108: 16428-16433, 2011.
- 853 **Fedorenko E, Duncan J, Kanwisher N.** Language-Selective and Domain-General
854 Regions Lie Side by Side within Broca's Area. *Curr Bio* 22: 2059-2062, 2012a.
- 855 **Fedorenko E, Nieto-Castañón A, Kanwisher N.** Lexical and syntactic representations
856 in the brain: An fMRI investigation with multi-voxel pattern analysis.
857 *Neuropsychologia* 50: 499-513, 2012b.
- 858 **Fedorenko E, McDermott JH, Norman-Haignere S, Kanwisher N.** Sensitivity to
859 musical structure in the human brain. *J Neurophysiol* 108: 3289-3300, 2012c.
- 860 **Fedorenko E, Duncan J, Kanwisher N.** Broad domain generality in focal regions of
861 frontal and parietal cortex. *P Natl Acad Sci USA* 110: 16616-16621, 2013.
- 862 **Fedorenko E.** The role of domain-general cognitive control in language comprehension.

- 863 *Front Psychol* 5:1-17, 2014.
- 864 **Fedorenko E, Thompson-Schill SL.** Reworking the language network. *Trends Cogn Sci*
865 18: 120-126, 2014.
- 866 **Fedorenko E, Varley R.** Language and thought are not the same thing: evidence from
867 neuroimaging and neurological patients. *Ann NY Acad Sci* 1369: 132-153, 2016.
- 868 **Fiebach CJ, Schubotz RI.** Dynamic anticipatory processing of hierarchical sequential
869 events: a common role for Broca's area and ventral premotor cortex across
870 domains?. *Cortex* 42: 499-502, 2006.
- 871 **Fischl B, Rajendran N, Busa E, Augustinack J, Hinds O, Yeo BTT, Mohlberg H,**
872 **Amunts K, Zilles K.** Cortical Folding Patterns and Predicting Cytoarchitecture.
873 *Cereb Cortex* 18: 1973-1980, 2008.
- 874 **Flash T, Bizzi E.** Cortical circuits and modules in movement generation: experiments
875 and theories. *Curr Opin Neurobiol* 41: 174-178, 2016.
- 876 **Flinker A, Korzeniewska A, Shestyuk AY, Franaszczuk PJ, Dronkers NF, Knight**
877 **RT, Crone NE.** Redefining the role of Broca's area in speech. *Proc Natl Acad Sci*
878 *USA* 112: 2871-2875, 2015.
- 879 **Frost MA, Goebel R.** Measuring structural-functional correspondence: Spatial
880 variability of specialised brain regions after macro-anatomical alignment.
881 *NeuroImage* 59: 1369-1381, 2012.
- 882 **Gallivan JP, Culham JC.** Neural coding within human brain areas involved in actions.
883 *Curr Opin Neurobiol* 33: 141-149, 2015.
- 884 **Goldin-Meadow S, Brentari D.** Gesture, sign, and language: The coming of age of sign
885 language and gesture studies. *Behav Brain Sci* 1-82, 2016.

- 886 **Guenther FH.** *Neural Control of Speech*. Cambridge, MA: MIT Press, 2016.
- 887 **Häberling IS, Corballis PM, Corballis MC.** Language, gesture, and handedness:
888 Evidence for independent lateralized networks. *Cortex* 82: 72-85, 2016.
- 889 **Harris KD, Shepherd GMG.** The neocortical circuit: themes and variations. *Nat*
890 *Neurosci* 16: 170-181, 2015.
- 891 **Hauk O, Johnsrude I, Pulvermüller F.** Somatotopic Representation of Action Words in
892 Human Motor and Premotor Cortex. *Neuron* 41: 301-307, 2004.
- 893 **Hickok G, Okada K, Serences JT.** Area Spt in the Human Planum Temporale Supports
894 Sensory-Motor Integration for Speech Processing. *J Neurophysiol* 101: 2725-
895 2732, 2009.
- 896 **Hickok G.** Eight Problems for the Mirror Neuron Theory of Action Understanding in
897 Monkeys and Humans. *J Cogn Neurosci* 21: 1229-1243, 2009.
- 898 **Higuchi S, Chaminade T, Imamizu H, Kawato M.** Shared neural correlates for
899 language and tool use in Broca's area. *NeuroReport* 20: 1376-1381, 2009.
- 900 **Hommel B, Müsseler J, Aschersleben G, Prinz W.** The Theory of Event Coding
901 (TEC): a framework for perception and action planning. *Behav Brain Sci* 24:
902 849-878, 2001.
- 903 **Hugdahl K, Raichle ME, Mitra A, Specht K.** On the existence of a generalized non-
904 specific task-dependent network. *Front Hum Neurosci* 9: 430, 2015.
- 905 **Johnson-Frey SH, Newman-Norlund R, Grafton ST.** A Distributed Left Hemisphere
906 Network Active During Planning of Everyday Tool Use Skills. *Cereb Cortex* 15:
907 681-695, 2005.
- 908 **Juch H, Zimine I, Seghier ML, Lazeyras F, Fasel JHD.** Anatomical variability of the

Action response in language regions

- 909 lateral frontal lobe surface: Implication for intersubject variability in language
910 neuroimaging. *NeuroImage* 24: 504-514, 2005.
- 911 **Julian JB, Fedorenko E, Webster J, Kanwisher N.** An algorithmic method for
912 functionally defining regions of interest in the ventral visual pathway.
913 *NeuroImage* 60: 2357-2364, 2012.
- 914 **Kimura D.** Acquisition of a motor skill after left-hemisphere damage. *Brain* 100: 527-
915 542, 1977.
- 916 **Kimura D, Battison R, Lubert B.** Impairment of Nonlinguistic Hand Movements in a
917 Deaf Aphasic. *Brain Lang* 3: 566-571, 1976.
- 918 **Koechlin E, Ody C, Kouneiher F.** The Architecture of Cognitive Control in the Human
919 Prefrontal Cortex. *Science* 302: 1181-1185, 2003.
- 920 **Koechlin E, Jubault T.** Broca's Area and the Hierarchical Organization of Human
921 Behavior. *Neuron* 50: 963-974, 2006.
- 922 **Koechlin E, Summerfield C.** An information theoretical approach to prefrontal
923 executive function. *Trends Cogn Sci* 11: 229-235, 2007.
- 924 **Koelsch S, Gunter TC, Cramon DY, Zysset S, Lohmann G, Friederici AD.** Bach
925 Speaks: A Cortical "Language-Network" Serves the Processing of Music.
926 *NeuroImage* 17: 956-966, 2002.
- 927 **Kriegeskorte N.** Pattern-information analysis: From stimulus decoding to computational-
928 model testing. *NeuroImage* 56: 411-421, 2011.
- 929 **Lambon Ralph MA, Jefferies E, Patterson K, Rogers TT.** The neural and
930 computational bases of semantic cognition. *Nature* 18: 42-55, 2017.
- 931 **Leshinskaya A, Caramazza A.** For a cognitive neuroscience of concepts: moving

- 932 beyond the grounding issue. *Psychon Bull Rev* 23: 991-1001, 2016.
- 933 **Lingnau A, Gesierich B, Caramazza A.** Asymmetric fMRI adaptation reveals no
934 evidence for mirror neurons in humans. *P Natl Acad Sci USA* 106: 9925-9930,
935 2009.
- 936 **Maess B, Koelsch S, Gunter TC, Friederici AD.** Musical syntax is processed in Broca's
937 area: an MEG study. *Nat Neurosci* 4: 540-545, 2001.
- 938 **Mahowald K, Fedorenko E.** Reliable individual-level neural markers of high-level
939 language processing: A necessary precursor for relating neural variability to
940 behavioral and genetic variability. *NeuroImage* 139: 74-93, 2016.
- 941 **Marstaller L, Burianová H.** The multisensory perception of co-speech gestures – A
942 review and meta-analysis of neuroimaging studies. *J Neurolinguist* 30: 69-77,
943 2014.
- 944 **McNeill D.** *Hand and Mind: What Gestures Reveal about Thought*. Chicago, OH:
945 University of Chicago Press, 1992.
- 946 **Meister IG, Iacoboni M.** No Language-Specific Activation during Linguistic Processing
947 of Observed Actions. *PLoS ONE* 2: e891, 2007.
- 948 **Menenti L, Gierhan SME, Segaert K, Hagoort P.** Shared Language: Overlap and
949 Segregation of the Neuronal Infrastructure for Speaking and Listening Revealed
950 by Functional MRI. *Psychol Sci* 22: 1173-1182, 2011.
- 951 **Meteyard L, Rodriguez Cuadrado S, Bahrami B, Vigliocco G.** Coming of age: a
952 review of embodiment and the neuroscience of semantics. *Cortex* 48: 788-804,
953 2012.
- 954 **Molenberghs P, Cunnington R, Mattingley JB.** Brain regions with mirror properties: A

Action response in language regions

- 955 meta-analysis of 125 human fMRI studies. *Neurosci Biobehav R* 36: 341-349,
956 2012.
- 957 **Monti MM, Osherson DN.** Logic, language and the brain. *Brain Res* 1428: 33-42, 2012.
- 958 **Mukamel R, Ekstrom AD, Kaplan J, Iacoboni M, Fried I.** Single-Neuron Responses
959 in Humans during Execution and Observation of Actions. *Curr Biol* 20: 750-756,
960 2010.
- 961 **Murakami T, Restle J, Ziemann U.** Observation-execution matching and action
962 inhibition in human primary motor cortex during viewing of speech-related lip
963 movements or listening to speech. *Neuropsychologia* 49: 2045-2054, 2011.
- 964 **Nieto-Castañón A, Fedorenko E.** Subject-specific functional localizers increase
965 sensitivity and functional resolution of multi-subject analyses. *NeuroImage* 63:
966 1646-1669, 2012.
- 967 **Norman-Haignere S, Kanwisher NG, McDermott JH.** Distinct Cortical Pathways for
968 Music and Speech Revealed by Hypothesis-Free Voxel Decomposition. *Neuron*
969 88: 1281-1296, 2015.
- 970 **Oldfield RC.** The Assessment and Analysis of Handedness: The Edinburgh Inventory.
971 *Neuropsychologia* 9: 97-113, 1971.
- 972 **Overath T, McDermott JH, Zarate JM, Poeppel D.** The cortical analysis of speech-
973 specific temporal structure revealed by responses to sound quilts. *Nat Neurosci*
974 18: 903-911, 2015.
- 975 **Papagno C, Della Sala S, Basso A.** Ideomotor apraxia without aphasia and aphasia
976 without apraxia: the anatomical support for a double dissociation. *J Neurol*
977 *Neurosurg Ps* 56: 286-289, 1993.

- 978 **Petrides M, Pandya DN.** Distinct Parietal and Temporal Pathways to the Homologues of
979 Broca's Area in the Monkey. *PLoS Biol* 7: e1000170, 2009.
- 980 **Pierrot-Deseilligny C, Milea D, Müri RM.** Eye movement control by the cerebral
981 cortex. *Curr Opin Neurol* 17: 17-25, 2004.
- 982 **Pischedda D, Görgen K, Haynes JD, Reverberi C.** Neural Representations of
983 Hierarchical Rule Sets: The Human Control System Represents Rules Irrespective
984 of the Hierarchical Level to Which They Belong. *J Neurosci* 37: 12281-12296,
985 2017.
- 986 **Pitcher D, Dilks DD, Saxe RR, Triantafyllou C, Kanwisher N.** Differential selectivity
987 for dynamic versus static information in face-selective cortical regions.
988 *NeuroImage* 56: 2356-2363, 2011.
- 989 **Poldrack RA.** Can cognitive processes be inferred from neuroimaging data?. *Trends*
990 *Cogn Sci* 10: 59-63, 2006.
- 991 **Poldrack RA.** Inferring Mental States from Neuroimaging Data: From Reverse Inference
992 to Large-Scale Decoding. *Neuron* 72: 692-697, 2011.
- 993 **Pulvermüller F, Huss M, Kherif F, Moscoso del Prado Martin F, Hauk O, Shtyrov**
994 **Y.** Motor cortex maps articulatory features of speech sounds. *Proc Nat Acad Sci*
995 *USA* 103: 7865-7870, 2006.
- 996 **Pulvermüller F, Fadiga L.** Active perception: sensorimotor circuits as a cortical basis
997 for language. *Nat Neuro* 11: 351-360, 2010.
- 998 **Redcay E, Ludlum RS, Velnoskey KR, Kanwal S.** Communicative Signals Promote
999 Object Recognition Memory and Modulate the Right Posterior STS. *J Cognitive*
1000 *Neurosci* 28: 8-19, 2016.

- 1001 **Rizzolatti G, Camarda R, Fogassi L, Gentilucci M, Luppino G, Matelli M.** Function
1002 organization of inferior area 6 in the macaque monkey. *Exp Brain Res* 71: 491-
1003 507, 1988.
- 1004 **Rizzolatti G, Arbib MA.** Language within our grasp. *Trends Neurosci* 21: 188-194,
1005 1998.
- 1006 **Rosenbaum DA, Meulenbroek RJ, Vaughan J, Jansen C.** Posture-Based Motion
1007 Planning: Applications to Grasping. *Psychol Rev* 108: 709-734, 2001.
- 1008 **Saxe R, Brett M, Kanwisher N.** Divide and conquer: A defense of functional localizers.
1009 *NeuroImage* 30: 1088-1096, 2006.
- 1010 **Saygin AP, Wilson SM, Dronkers NF, Bates E.** Action comprehension in aphasia:
1011 Linguistic and non-linguistic deficits and their lesion correlates.
1012 *Neuropsychologia* 42: 1788-1804, 2004.
- 1013 **Schack T.** The cognitive architecture of complex movement. *Int J Sport Exercise Psych*
1014 2: 403-438, 2004.
- 1015 **Scott TL, Gallée J, Fedorenko E.** A new fun and robust version of an fMRI localizer for
1016 the frontotemporal language system. *Cogn Neurosci* 167-176, 2016.
- 1017 **Shepherd SV, Freiwald WA.** Functional Networks for Social Communication in the
1018 Macaque Monkey. *Neuron* 99: 413-420, 2018.
- 1019 **Simmons WK, Ramjee V, Beauchamp MS, McRae K, Martin A, Barsalou LW.** A
1020 common neural substrate for perceiving and knowing about color.
1021 *Neuropsychologia* 45: 2802-2810, 2007.
- 1022 **Sirigu A, Cohen L, Zalla T, Pradat-Diehl P, Van Eeckhout P, Grafman J, Agid Y.**
1023 Distinct Frontal Regions for Processing Sentence Syntax and Story Grammar.

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- 1024 *Cortex* 34: 771-778, 1998.
- 1025 **Slocombe KE.** Vocal Communication in Primates. In: *Emerging Trends in the Social and*
1026 *Behavioral Sciences: An Interdisciplinary, Searchable, and Linkable Resource*,
1027 edited by Kosslyn SM. Hoboken, NJ: John Wiley & Sons, Inc, 2015, p. 1-12.
- 1028 **Tahmasebi AM, Davis MH, Wild CJ, Rodd JM, Hakyemez H, Abolmaesumi P,**
1029 **Johnsrude IS.** Is the Link between Anatomical Structure and Function Equally
1030 Strong at All Cognitive Levels of Processing?. *Cereb Cortex* 22: 1593-1603, 2011.
- 1031 **Tarhan L, Konkle T.** Modeling the Neural Structure Underlying Human Action
1032 Perception [Online]. Conference on Cognitive Computational Neuroscience.
1033 <https://www2.securecms.com/CCNeuro/docs-0/591b2fa768ed3fa74b57b899.pdf>
1034 [2017].
- 1035 **Tettamanti M, Weniger D.** Broca's area: A supramodal hierarchical processor?. *Cortex*
1036 42: 491-494, 2006.
- 1037 **Thesen S, Heid O, Mueller E, Schad L.** Prospective Acquisition Correction for Head
1038 Motion With Image-Based Tracking for Real-Time fMRI. *Magn Reson Med* 44:
1039 457-465, 2000.
- 1040 **Tomaiuolo F, MacDonald JD, Caramanos Z, Posner G, Chiavaras M, Evans AC,**
1041 **Petrides M.** Morphology, morphometry and probability mapping of the pars
1042 opercularis of the inferior frontal gyrus: an in vivo MRI analysis. *Eur J Neurosci*
1043 11: 3033-3046, 1999.
- 1044 **Tomasello M.** *Origins of Human Communication*. Cambridge, MA: MIT Press, 2008.
- 1045 **Tranel D, Kemmerer D, Adolphs R, Damasio H, Damasio AR.** Neural correlates of
1046 conceptual knowledge for actions. *Cogn Neuropsychol* 20: 409-432, 2003.

- 1047 **Tremblay P, Dick AS.** Broca and Wernicke are dead, or moving past the classic model
1048 of language neurobiology. *Brain Lang* 162: 60-71, 2016.
- 1049 **Tzourio-Mazoyer N, Landeau B, Papathanassiou D, Crivello F, Etard O, Delcroix**
1050 **N, Mazoyer B, Joliot M.** Automated Anatomical Labeling of Activations in SPM
1051 Using a Macroscopic Anatomical Parcellation of the MNI MRI Single-Subject
1052 Brain. *NeuroImage* 15: 273-289, 2002.
- 1053 **Villareal M, Fridman EA, Amengual A, Falasco G, Gercovich ER, Ulloa ER,**
1054 **Leiguarda RC.** The neural substrate of gesture recognition. *Neuropsychologia*
1055 46: 2371-2382, 2008.
- 1056 **Watkins KE, Strafella AP, Paus T.** Seeing and hearing speech excites the motor system
1057 involved in speech production. *Neuropsychologia* 41: 989-994, 2003.
- 1058 **Willems RM, Hagoort P.** Neural evidence for the interplay between language, gesture,
1059 and action: A review. *Brain Lang* 101: 278-289, 2007.
- 1060 **Willems RM, Van der Haegen L, Fisher SE, Francks C.** On the other hand: including
1061 left-handers in cognitive neuroscience and neurogenetics. *Nat Neuro* 15: 193-201,
1062 2014.
- 1063 **Xu J, Gannon PJ, Emmorey K, Smith JF, Braun AR.** Symbolic gestures and spoken
1064 language are processed by a common neural system. *P Natl Acad Sci USA* 106:
1065 20664-20669, 2009.
- 1066 **Yang J, Andric M, Mathew MM.** The neural basis of hand gesture comprehension: A
1067 meta-analysis of functional magnetic resonance imaging studies. *Neurosci*
1068 *Biobehav R* 57: 88-104, 2015.
- 1069
1070

1071 **TABLES**1072 **Table 1.** Timing parameters for the different versions of the language localizer task.

	Version		
	A	B	C
Number of participants	5	5	80
Task: Passive Reading or Memory?	M	M	PR
Words / nonwords per trial	8	12	12
Trial duration (ms)	4,800	6,000	6,000
Fixation	300	300	100
Presentation of each word / nonword	350	350	450
Fixation	---	---	500
Memory probe	1,350	1,000	---
Fixation	350	500	---
Trials per block	5	3	3
Block duration (s)	24	18	18
Blocks per condition (per run)	8	8	8
Conditions	Sentences	Sentences	Sentences
	Nonwords	Nonwords	Nonwords
Fixation block duration (s)	16	18	14
Number of fixation blocks	5	5	5
Total run time (s)	464	378	358
Number of runs	2	2	2

1073

1074 **Table 2.** Results for each experiment and condition for the six language fROIs. Note that
 1075 although in Figure 2, we plot the language localizer responses across the entire set of
 1076 participants in the current study, all the comparisons between the action conditions and
 1077 the conditions of the language localizer experiment were performed within each
 1078 experiment separately using two-tailed paired-samples *t*-tests. In columns 2 and 3,
 1079 significance values indicate whether the action observation/imitation condition elicited a
 1080 response reliably above the baseline (column 2) or reliably above the nonword condition
 1081 (column 3). In column 4, significance values indicate whether the action
 1082 observation/imitation condition elicited a response reliably below the sentence condition.

1083

Experiment	Condition	ROI	Action condition vs. fixation	Action condition vs. nonwords	Action condition vs. sentences
Expt. 1	hand action observation (attention to action)	<i>LIFGorb</i>	$t(9) = -2.29, p = 0.143$	$t(9) = -1.33, p = 0.357$	$t(9) = 4.18, p < 0.005$
		<i>LIFG</i>	$t(9) < 1 , n.s.$	$t(9) < 1 , n.s.$	$t(9) = 3.31, p < 0.05$
		<i>LMFG</i>	$t(9) = 1.59, p = 0.294$	$t(9) = -1.66, p = 0.357$	$t(9) = 5.67, p < 0.005$
		<i>LAntTemp</i>	$t(9) = -3.32, p = 0.053$	$t(9) < 1 , n.s.$	$t(9) = 3.88, p < 0.01$
		<i>LPostTemp</i>	$t(9) < 1 , n.s.$	$t(9) = -1.26, p = 0.357$	$t(9) = 4.65, p < 0.005$
		<i>LangG</i>	$t(9) = -1.08, p = 0.463$	$t(9) = 2.33, p = 0.271$	$t(9) = 2.76, p < 0.05$
	hand action observation (attention to object)	<i>LIFGorb</i>	$t(9) = -4.35, p < 0.01$	$t(9) = -2.10, p = 0.130$	$t(9) = 5.42, p < 0.001$
		<i>LIFG</i>	$t(9) < 1 , n.s.$	$t(9) = -1.41, p = 0.288$	$t(9) = 4.24, p < 0.005$
		<i>LMFG</i>	$t(9) = 1.54, p = 0.234$	$t(9) = -2.45, p = 0.110$	$t(9) = 5.82, p < 0.001$
		<i>LAntTemp</i>	$t(9) = -5.82, p < 0.005$	$t(9) = -1.16, p = 0.291$	$t(9) = 4.72, p < 0.005$
		<i>LPostTemp</i>	$t(9) = -1.40, p = 0.234$	$t(9) = -2.52, p = 0.110$	$t(9) = 6.10, p < 0.001$

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		<i>LangG</i>	$t(9) = -2.87, p < 0.05$	$t(9) = 1.12, p = 0.291$	$t(9) = 4.75, p < 0.005$
Expt. 2	Face action observation	<i>LIFGorb</i>	$t(53) = 3.30, p < 0.005$	$t(53) < 1 , n.s.$	$t(53) = 6.19, p < 0.0001$
		<i>LIFG</i>	$t(53) = 3.28, p < 0.005$	$t(53) < 1 , n.s.$	$t(53) = 7.98, p < 0.0001$
		<i>LMFG</i>	$t(53) = 2.12, p < 0.05$	$t(53) = -3.48, p < 0.005$	$t(53) = 9.36, p < 0.0001$
		<i>LAntTemp</i>	$t(53) = 3.41, p < 0.005$	$t(53) = 1.66, p = 0.157$	$t(53) = 8.89, p < 0.0001$
		<i>LPostTemp</i>	$t(53) = 4.14, p < 0.0005$	$t(53) = -1.65, p = 0.157$	$t(53) = 9.69, p < 0.0001$
		<i>LangG</i>	$t(53) = 4.98, p < 0.0001$	$t(53) = 4.52, p < 0.0005$	$t(53) < 1 , n.s.$
	Body action observation	<i>LIFGorb</i>	$t(53) = 4.16, p < 0.0005$	$t(53) < 1 , n.s.$	$t(53) = 5.73, p < 0.0001$
		<i>LIFG</i>	$t(53) = 3.56, p < 0.005$	$t(53) < 1 , n.s.$	$t(53) = 7.76, p < 0.0001$
		<i>LMFG</i>	$t(53) = 3.46, p < 0.005$	$t(53) = -3.57, p < 0.005$	$t(53) = 10.50, p < 0.0001$
		<i>LAntTemp</i>	$t(53) = 2.87, p < 0.01$	$t(53) < 1 , n.s.$	$t(53) = 9.64, p < 0.0001$
		<i>LPostTemp</i>	$t(53) = 3.23, p < 0.005$	$t(53) = -2.22, p = 0.061$	$t(53) = 10.31, p < 0.0001$
		<i>LangG</i>	$t(53) = 6.66, p < 0.0001$	$t(53) = 6.00, p < 0.0001$	$t(53) = -1.83, p = 0.073$
Expt. 3a	Face action observation	<i>LIFGorb</i>	$t(12) = 1.76, p = 0.156$	$t(12) < 1 , n.s.$	$t(12) = 4.74, p < 0.001$
		<i>LIFG</i>	$t(12) = 2.53, p = 0.146$	$t(12) < 1 , n.s.$	$t(12) = 3.31, p < 0.01$
		<i>LMFG</i>	$t(12) < 1 , n.s.$	$t(12) = -2.11, p = 0.169$	$t(12) = 5.17, p < 0.001$
		<i>LAntTemp</i>	$t(12) < 1 , n.s.$	$t(12) < 1 , n.s.$	$t(12) = 5.24, p < 0.001$
		<i>LPostTemp</i>	$t(12) = 1.97, p = 0.146$	$t(12) = -1.10, p = 0.582$	$t(12) = 4.78, p < 0.001$
		<i>LangG</i>	$t(12) = 2.19, p = 0.146$	$t(12) = 2.65, p = 0.126$	$t(12) = 1.13, p = 0.282$
	Eye action observation	<i>LIFGorb</i>	$t(12) < 1 , n.s.$	$t(12) < 1 , n.s.$	$t(12) = 4.03, p < 0.005$
		<i>LIFG</i>	$t(12) = 2.25, p = 0.263$	$t(12) < 1 , n.s.$	$t(12) = 3.64, p < 0.005$
		<i>LMFG</i>	$t(12) = 1.40, p = 0.562$	$t(12) = -1.20, p = 0.509$	$t(12) = 4.33, p < 0.005$
		<i>LAntTemp</i>	$t(12) < 1 , n.s.$	$t(12) < 1 , n.s.$	$t(12) = 6.95, p < 0.0001$
		<i>LPostTemp</i>	$t(12) < 1 , n.s.$	$t(12) = -1.37, p = 0.509$	$t(12) = 5.20, p < 0.001$
		<i>LangG</i>	$t(12) < 1 , n.s.$	$t(12) = 1.66, p = 0.509$	$t(12) = 1.62, p = 0.131$
	Mouth action observation	<i>LIFGorb</i>	$t(12) < 1 , n.s.$	$t(12) = -2.11, p = 0.113$	$t(12) = 5.72, p < 0.0005$
		<i>LIFG</i>	$t(12) = 1.29, p = 0.440$	$t(12) < 1 , n.s.$	$t(12) = 4.10, p < 0.005$
		<i>LMFG</i>	$t(12) < 1 , n.s.$	$t(12) = -2.54, p = 0.078$	$t(12) = 5.11, p < 0.001$
		<i>LAntTemp</i>	$t(12) < 1 , n.s.$	$t(12) < 1 , n.s.$	$t(12) = 6.93, p < 0.0001$
		<i>LPostTemp</i>	$t(12) = 1.77, p = 0.328$	$t(12) = -1.20, p = 0.383$	$t(12) = 4.69, p < 0.001$
		<i>LangG</i>	$t(12) = 1.73, p = 0.328$	$t(12) = 3.68, p < 0.05$	$t(12) = 1.04, p = 0.318$
	Hand action observation	<i>LIFGorb</i>	$t(12) < 1 , n.s.$	$t(12) = -1.14, p = 0.353$	$t(12) = 4.80, p < 0.001$
		<i>LIFG</i>	$t(12) = 2.04, p = 0.127$	$t(12) < 1 , n.s.$	$t(12) = 4.24, p < 0.005$
		<i>LMFG</i>	$t(12) = 2.83, p < 0.05$	$t(12) = -1.81, p = 0.190$	$t(12) = 5.13, p < 0.0005$
		<i>LAntTemp</i>	$t(12) = -1.55, p = 0.220$	$t(12) = -1.10, p = 0.353$	$t(12) = 5.84, p < 0.0005$
		<i>LPostTemp</i>	$t(12) = 1.15, p = 0.326$	$t(12) = -2.92, p < 0.05$	$t(12) = 6.48, p < 0.0005$
		<i>LangG</i>	$t(12) = 3.31, p < 0.05$	$t(12) = 4.38, p < 0.01$	$t(12) < 1 , n.s.$
Expt. 3b	Face action imitation	<i>LIFGorb</i>	$t(12) = 2.06, p = 0.123$	$t(12) < 1 , n.s.$	$t(12) = 2.06, p = 0.075$
		<i>LIFG</i>	$t(12) = 1.08, p = 0.455$	$t(12) < 1 , n.s.$	$t(12) = 3.27, p < 0.05$
		<i>LMFG</i>	$t(12) = 3.09, p = 0.056$	$t(12) = -2.11, p = 0.169$	$t(12) = 2.52, p < 0.05$
		<i>LAntTemp</i>	$t(12) < 1 , n.s.$	$t(12) < 1 , n.s.$	$t(12) = 4.62, p < 0.005$
		<i>LPostTemp</i>	$t(12) = 2.37, p = 0.107$	$t(12) = -1.10, p = 0.582$	$t(12) = 4.10, p < 0.005$
		<i>LangG</i>	$t(12) < 1 , n.s.$	$t(12) = 2.65, p = 0.126$	$t(12) = 1.77, p = 0.101$
	Eye action imitation	<i>LIFGorb</i>	$t(12) = 1.26, p = 0.386$	$t(12) < 1 , n.s.$	$t(12) = 1.78, p = 0.101$
		<i>LIFG</i>	$t(12) = 1.03, p = 0.386$	$t(12) < 1 , n.s.$	$t(12) = 3.17, p < 0.05$
		<i>LMFG</i>	$t(12) = 3.39, p < 0.05$	$t(12) = -1.47, p = 0.335$	$t(12) = 2.25, p = 0.053$
		<i>LAntTemp</i>	$t(12) = -1.04, p = 0.386$	$t(12) = -2.17, p = 0.304$	$t(12) = 6.06, p < 0.0005$
		<i>LPostTemp</i>	$t(12) = 1.44, p = 0.386$	$t(12) = -1.7, p = 0.335$	$t(12) = 5.16, p < 0.001$
		<i>LangG</i>	$t(12) < 1 , n.s.$	$t(12) < 1 , n.s.$	$t(12) = 3.23, p < 0.05$
	Mouth action imitation	<i>LIFGorb</i>	$t(12) < 1 , n.s.$	$t(12) < 1 , n.s.$	$t(12) = 2.84, p < 0.05$
		<i>LIFG</i>	$t(12) = 1.27, p = 0.274$	$t(12) < 1 , n.s.$	$t(12) = 3.01, p < 0.05$
		<i>LMFG</i>	$t(12) = 3.24, p < 0.05$	$t(12) < 1 , n.s.$	$t(12) = 2.52, p < 0.05$
		<i>LAntTemp</i>	$t(12) = 2.04, p = 0.144$	$t(12) < 1 , n.s.$	$t(12) = 7.24, p < 0.0001$

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Hand action imitation	<i>LPostTemp</i>	$t(12) = 1.85, p = 0.144$	$t(12) < 1 , n.s.$	$t(12) = 5.10, p < 0.001$
	<i>LAngG</i>	$t(12) = 1.81, p = 0.144$	$t(12) = 1.89, p = 0.495$	$t(12) = 1.22, p = 0.247$
	<i>LIFGorb</i>	$t(12) = 1.32, p = 0.319$	$t(12) < 1 , n.s.$	$t(12) = 1.52, p = 0.163$
	<i>LIFG</i>	$t(12) = 1.78, p = 0.263$	$t(12) = 1.15, p = 0.549$	$t(12) = 1.77, p = 0.163$
	<i>LMFG</i>	$t(12) = 3.80, p < 0.05$	$t(12) = 1.34, p = 0.549$	$t(12) = 1.49, p = 0.163$
	<i>LAntTemp</i>	$t(12) < 1 , n.s.$	$t(12) < 1 , n.s.$	$t(12) = 4.76, p < 0.005$
	<i>LPostTemp</i>	$t(12) = 1.62, p = 0.263$	$t(12) < 1 , n.s.$	$t(12) = 4.16, p < 0.005$
	<i>LAngG</i>	$t(12) < 1 , n.s.$	$t(12) = 1.23, p = 0.549$	$t(12) = 1.49, p = 0.163$

Table 3. Results for each experiment (for Experiments 2 and 3a/b; no MD localizer was included in Experiment 1) and condition for the six MD fROIs. Significance values indicate whether the action observation/imitation condition elicited a response reliably above the baseline.

Experiment	Condition	ROI	Action condition vs. fixation
Expt. 2	Face action observation	<i>LIFGop</i>	$t(54) < 1 , n.s.$
		<i>RIFGop</i>	$t(54) = 1.15, p = 0.521$
		<i>LPrecG</i>	$t(54) < 1 , n.s.$
		<i>RPrecG</i>	$t(54) = 2.25, p = 0.169$
		<i>LParSup</i>	$t(54) = -1.66, p = 0.449$
		<i>RParSup</i>	$t(54) = -1.28, p = 0.521$
	Body action observation	<i>LIFGop</i>	$t(54) = 1.58, p = 0.197$
		<i>RIFGop</i>	$t(54) = 1.89, p = 0.521$
		<i>LPrecG</i>	$t(54) = 3.06, p < 0.05$
		<i>RPrecG</i>	$t(54) = 4.51, p < 0.0005$
		<i>LParSup</i>	$t(54) = 4.82, p < 0.0005$
		<i>RParSup</i>	$t(54) = 4.22, p < 0.001$
Expt. 3a	Face action observation	<i>LIFGop</i>	$t(12) = 1.27, p = 0.516$
		<i>RIFGop</i>	$t(12) = 1.46, p = 0.514$
		<i>LPrecG</i>	$t(12) = 1.40, p = 0.514$
		<i>RPrecG</i>	$t(12) = 1.97, p = 0.514$
		<i>LParSup</i>	$t(12) = 1.36, p = 0.514$
		<i>RParSup</i>	$t(12) < 1 , n.s.$
	Eye action observation	<i>LIFGop</i>	$t(12) = 2.52, p = 0.081$
		<i>RIFGop</i>	$t(12) = 3.68, p < 0.05$
		<i>LPrecG</i>	$t(12) = 3.40, p < 0.05$
		<i>RPrecG</i>	$t(12) = 4.66, p < 0.01$
		<i>LParSup</i>	$t(12) = 3.15, p < 0.05$
		<i>RParSup</i>	$t(12) = 2.03, p = 0.141$
	Mouth action observation	<i>LIFGop</i>	$t(12) = 2.32, p = 0.349$
		<i>RIFGop</i>	$t(12) = 1.40, p = 0.507$
		<i>LPrecG</i>	$t(12) = 1.89, p = 0.499$
		<i>RPrecG</i>	$t(12) = 2.47, p = 0.349$
		<i>LParSup</i>	$t(12) = 1.37, p = 0.507$
		<i>RParSup</i>	$t(12) < 1 , n.s.$
	Hand action observation	<i>LIFGop</i>	$t(12) = 2.00, p = 0.207$
		<i>RIFGop</i>	$t(12) = 2.38, p = 0.126$
		<i>LPrecG</i>	$t(12) = 2.67, p = 0.091$
		<i>RPrecG</i>	$t(12) = 3.91, p < 0.05$
		<i>LParSup</i>	$t(12) = 2.80, p = 0.091$
		<i>RParSup</i>	$t(12) = 1.83, p = 0.236$

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Expt. 3b	Face action imitation	<i>LIFGop</i>	$t(12) = 2.87, p < 0.05$
		<i>RIFGop</i>	$t(12) = 3.23, p < 0.05$
		<i>LPrecG</i>	$t(12) = 4.50, p < 0.005$
		<i>RPrecG</i>	$t(12) = 7.56, p < 0.0005$
		<i>LParSup</i>	$t(12) = 5.23, p < 0.001$
		<i>RParSup</i>	$t(12) = 3.29, p < 0.05$
	Eye action imitation	<i>LIFGop</i>	$t(12) = 2.09, p = 0.117$
		<i>RIFGop</i>	$t(12) = 2.71, p < 0.05$
		<i>LPrecG</i>	$t(12) = 2.78, p < 0.05$
		<i>RPrecG</i>	$t(12) = 3.64, p < 0.05$
		<i>LParSup</i>	$t(12) = 3.05, p < 0.05$
		<i>RParSup</i>	$t(12) = 1.80, p = 0.164$
	Mouth action imitation	<i>LIFGop</i>	$t(12) = 3.97, p < 0.01$
		<i>RIFGop</i>	$t(12) = 3.26, p < 0.05$
		<i>LPrecG</i>	$t(12) = 4.43, p < 0.005$
		<i>RPrecG</i>	$t(12) = 4.69, p < 0.005$
		<i>LParSup</i>	$t(12) = 4.17, p < 0.005$
		<i>RParSup</i>	$t(12) = 2.05, p = 0.088$
	Hand action imitation	<i>LIFGop</i>	$t(12) = 3.38, p < 0.01$
		<i>RIFGop</i>	$t(12) = 4.44, p < 0.005$
		<i>LPrecG</i>	$t(12) = 4.50, p < 0.005$
		<i>RPrecG</i>	$t(12) = 4.83, p < 0.005$
		<i>LParSup</i>	$t(12) = 4.24, p < 0.005$
		<i>RParSup</i>	$t(12) = 4.50, p < 0.005$

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Figure legends

Figure 1: Sample stimuli for each experiment and condition. *Experiment 1.* a. Example objects, grouped vertically by family. b. Example family of dissimilar actions. *Experiment 2.* c. Example body action stimuli. d. Two sample face action stimuli. *Experiment 3a/b.* e. Example face actions. f. Example eye actions. g. Example mouth actions. h. Example hand actions.

Figure 2: Response to the language localizer conditions (estimated in data not used for fROI definition, as described in [Methods](#)) and action conditions across experiments. Next to each bar graph, we show the language parcels used to constrain the selection of individual language fROIs; the individual fROIs constitute 10% of each parcel (see [Methods](#) for details). Error bars indicate standard errors of the mean over participants.

Figure 3: Responses in multiple-demand regions to the action conditions in Experiments 2 and 3a/b. Next to each bar graph, we show the MD parcels used to constrain the selection of individual MD fROIs; the individual fROIs constitute 10% of each parcel (see [Methods](#) for details). Error bars indicate standard errors of the mean over participants.

Figure 4: a. Responses in speech-responsive functional regions of interest (fROIs) in the auditory cortex (defined by nonword perception > hand action observation contrast in Experiment 3a, see [Methods](#) for details) to the nonword condition and the four action observation conditions in Experiment 3a. Responses are estimated using data not used for

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1120 fROI definition (see Methods). Error bars indicate standard error of the mean over
1121 participants. Abbreviations, from left to right: L/R PP – left/right planum polare, L/R PT
1122 – left/right planum temporale, L/R ASTG – left/right anterior superior temporal gyrus,
1123 L/R PSTG – left/right posterior superior temporal gyrus.

1124 b. Responses in articulation-responsive fROIs in the premotor cortex (defined by the
1125 nonword imitation > hand action imitation contrast in Experiment 3b, see Methods for
1126 details) to the nonword imitation condition and the four action imitation conditions in
1127 Experiment 3b. Responses are estimated using data not used for fROI definition (see
1128 Methods). Error bars indicate standard error of the mean over participants. Abbreviations,
1129 from left to right: LPrCG – left precentral gyrus, RiPrCG – right inferior precentral gyrus,
1130 RsPrCG – right superior precentral gyrus.

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